510(K)Summary

"This summary of 510(k) safety and effectiveness information is being submitted in accordance with the requirements of SMDA 1990 and 21 CFR 807.92."

"The assigned 510(k) number is: KIDDL6" (applicant leave blank)

Premarket Notification [510(k)] Summary

[(a)(1)]. The summary contains on the first page, preferably on your letterhead paper, the submitter's name, address, phone and fax numbers, name of contact person, and date the summary was prepared:

Submitter's name:

SHANDONG JUNFENG INDUSTRIES CO., LTD

Submitter's address:

No.899, Hengtong Road, Zhoucun, Zibo, Shandong,

255300, China

Phone number:

(86) 533-6537122

Fax number:

(86) 533-6537122

Name of contact person:

Ms. BIAN Qinyu

Date the summary was prepared:

Jul. 04th, 2012

[(a)(2)]. The name of the device, including the trade or proprietary name if applicable, the common or usual name, and the classification name, if known

Device Name:

Powder Free Vinyl Patient Examination Gloves, Clear (non-

colored)

Proprietary/Trade name:

Powder Free Vinyl Patient Examination Gloves

Common Name:

Patient examination glove

Classification Name:

Patient examination glove

Device Classification:

I

Regulation Number:

21 CFR 880.6250

Panel:

General Hospital (80)

Product Code:

LYZ

[(a)(3)]. An identification of the legally marketed device to which your firm is claiming substantial equivalence.

Class I* Powder Free Vinyl Patient Examination Gloves, Clear (non-colored) that meets all of the requirements of ASTM standard D 5250-06.

Predicate device: FUGUAN (Brand) Powder-Free Vinyl Patient Examination Gloves, Shijiazhuang Fuguan Plastic Products Co., Ltd., K032908.

[(a)(4)] A description of the device

Device Description: Powder Free Vinyl Patient Examination Gloves, Clear (non-colored)that meets all of the requirements of ASTM standard D 5250-06.

[(a)(5)] The summary describes the intended use of the device

Device Intended Use: Powder Free Vinyl Patient Examination Gloves, Clear (non-colored) is a disposable device intended for medical purposes that is worn on the examiner's hand or finger to prevent contamination between patient and examiner.

[(a)(6)] A summary of the technological characteristics of new device compared to the predicate device.

The powder free vinyl patient examination gloves, non sterile are summarized with the following technological characteristics compared to ASTM or equivalent standard.

Characteristics	Standard	Device performance
Dimension	ASTM standard D 5250-06	Meets
Physical Properties	ASTM standard D 5250-06	Meets
Freedom from pinholes	21 CFR 800.20	Meets
Powder Residual	ASTM standard D 5250-06	Meets
	and D6124-06	<2mg/glove
Biocompatability	Primary Skin Irritation in rabbits	Passes
•		Not a Primary Skin Irritation
	Dermal sensitization in the guinea pig	Passes
		Not a Dermal sensitization

[(b)(1)] A brief discussion of the non-clinical testing data that was submitted, referenced or relied on to demonstrate that the Subject Device is safe and effective, and whose performance meets the requirements of its pre-defined acceptance criteria and intended uses.

Powder Free Vinyl Patient Examination Gloves, Clear (non-colored)meet requirements per ASTM D5250-06, per ASTM D6124-06, per 21 CFR 800.20 and ISO10993-10. , it is safe and effective, and it's performance meets the requirements of its pre-defined acceptance criteria and intended uses.

[(b)(2)] A brief discussion of the clinical submitted, reference, or relied on in the premarket notification submission for a determination of substantial equivalence.

Clinical data is not needed for gloves or for most devices cleared by the 510(k) process.

[(b)(3)] The conclusions drawn from the nonclinical and clinical tests that demonstrate that the device is as safe, as effective, and performed as well or better than the legally marketed device identified in (a)(3).

It can be concluded that the Powder Free Vinyl Patient Examination Gloves meet the ASTM standard or equivalent standard and FDA requirements for waterleak test on pinhole AQL., meet labeling claims. It is as safe, as effective, and performed as well the legally marketed device identified in (a)(3).

Section 15

Substantial Equivalence Discussion

The Powder Free Vinyl Patient Examination Gloves, Clear (non-colored) made by SHANDONG JUNFENG INDUSTRIES CO., LTD, have the same intended use and technological characteristics as the Predicate devices, FUGUAN (Brand) Powder-Free Vinyl Patient Examination Gloves made by Shijiazhuang Fuguan Plastic Products Co., Ltd. K032908. The Subject Device is substantially equivalent to the predicate devices with regard to technologies, in design and very similar construction, material, function, and application. More detail information on the Table 1.

Based on the information in following Table I, the Subject Device (Powder Free Vinyl Patient Examination Gloves, Clear (non-colored)) has the same technological characteristics as the predicate device. The Powder Free Vinyl Patient Examination Gloves, Clear (non-colored) made by SHANDONG JUNFENG INDUSTRIES CO., LTD, are substantially equivalent to the predicate devices.

reatures & Description	Predicate Device	Medical Glove Guidance Manual(1661)	Subject Device	Result of Comparison
Intend for use	Powder free vinyl	Powder-Free	Powder-Free vinyl patient	
	patient examination	Examination Gloves:	examination gloves is a	
	glove is a disposable	A powder-free patient	disposable device	
	device intended for	examination glove is a	intended for medical	
	medical purposes	disposable device	purposes that is worn on	
	that is worn on the	intended for medical	the examiner's hand or	Substantially equivalent
	examiner's hand or	purposes that is worn on	finger to prevent	
	finger to prevent	the examiner's hand or	contamination between	
	contamination	finger to prevent	patient and examiner.	
	between patient and	contamination between		
	examiner.	patient and examiner.		
Device Description and		If viny1:		
Specifications		Do the vinyl examination		
		gloves meet all the		
	Mooth ACTM DEDEN	current specifications	Meets ASTM D5250	Substantially conjugat
	DOZOG INI DOZ GIBINI	listed under ASTM		oubstantianiy equivalent
		Specification D5250 or		
		an equivalent consensus		
		standard?		
Compare all materials	-	If the glove is made of a		
used to fabricate the	J/6	polymer or other type of	D/(C	Substantially equivalent
devices) }	material. identify the)	
		material.		

Dusting or Donning	PU	If a donning lubricant is	P.O.	Substantially equivalent
Powder:		used, state the		
		composition and		
		include biocompatibility		
		data for the lubricant in	-	
	·	an identified attachment;		
		also state the	-	
		name, manufacturer,		
		and address below		
	PU	Lubricant	Surface Coating Agent	Substantially equivalent
		'Generic Name/	PU-120-C	
		Lubricant		
		Brand Name		
Compare product	Meets ASTM D5250	We recommend you	Meets ASTM D5250	Substantially equivalent
specifications		certify that your finished	•	
		"powder-free" gloves		
		meet the following:		
		ASTM D 5250 standard		
		or an equivalent		
		standard for vinyl		
		A 4 4 L 1 L 1 L 1 L 1 L 1 L 1 L 1 L 1 L 1	Mooth	Substantially equivalent
Compare performance	Meets	At this time FDA		מתספומים לי הליים המיים
data supporting	ASTM D5151	recognizes the tollowing	ASTM Datat	
substantial equivalence	ASTM D5250	standards:	ASTM D5250	
-	ASTM D6124	Patient Examination	ASTM D6124	
		Gloves		
		ASTM D5151(Detection		- 13
		of Holes in Medical		

DEPARTMENT OF HEALTH & HUMAN SERVICES





Food and Drug Administration 10903 New Hampshire Avenue Document Control Room – WO66-G609 Silver Spring, MD 20993-0002

Shandong Junfeng Industries Company, Limited Mr. Chu Xiaoan Room 1606 Building 1, Jianxiang Yuan No. 209 Bei SI Huan Zhong Road Haidian District Beijing 100083, P.R. China SEP 2 7 2012

Re: K122266

Trade/Device Name: Powder-Free Vinyl Patient Examination Gloves,

Clear (Non-Colored)

Regulation Number: 21 CFR 880.6250

Regulation Name: Patient Examination Glove

Regulatory Class: I Product Code: LYZ Dated: September 6, 2012 Received: September 13, 2012

Dear Mr. Xiaoan:

We have reviewed your Section 510(k) premarket notification of intent to market the device referenced above and have determined the device is substantially equivalent (for the indications for use stated in the enclosure) to legally marketed predicate devices marketed in interstate commerce prior to May 28, 1976, the enactment date of the Medical Device Amendments, or to devices that have been reclassified in accordance with the provisions of the Federal Food, Drug, and Cosmetic Act (Act) that do not require approval of a premarket approval application (PMA). You may, therefore, market the device, subject to the general controls provisions of the Act include requirements for annual registration, listing of devices, good manufacturing practice, labeling, and prohibitions against misbranding and adulteration. Please note: CDRH does not evaluate information related to contract liability warranties. We remind you, however, that device labeling must be truthful and not misleading.

If your device is classified (see above) into either class II (Special Controls) or class III (PMA), it may be subject to additional controls. Existing major regulations affecting your device can be found in the Code of Federal Regulations, Title 21, Parts 800 to 898. In addition, FDA may publish further announcements concerning your device in the <u>Federal Register</u>.

Please be advised that FDA's issuance of a substantial equivalence determination does not mean that FDA has made a determination that your device complies with other requirements of the Act or any Federal statutes and regulations administered by other Federal agencies. You must comply with all the Act's requirements, including, but not limited to: registration and listing (21 CFR Part 807); labeling (21 CFR Part 801); medical device reporting (reporting of medical device-related adverse events) (21 CFR 803); good manufacturing practice requirements as set forth in the quality systems (QS) regulation (21 CFR Part 820); and if applicable, the electronic product radiation control provisions (Sections 531-542 of the Act); 21 CFR 1000-1050.

If you desire specific advice for your device on our labeling regulation (21 CFR Part 801), please go to

http://www.fda.gov/AboutFDA/CentersOffices/CDRH/CDRHOffices/ucm115809.htm for the Center for Devices and Radiological Health's (CDRH's) Office of Compliance. Also, please note the regulation entitled, "Misbranding by reference to premarket notification" (21CFR Part 807.97). For questions regarding the reporting of adverse events under the MDR regulation (21 CFR Part 803), please go to

http://www.fda.gov/MedicalDevices/Safety/ReportaProblem/default.htm for the CDRH's Office of Surveillance and Biometrics/Division of Postmarket Surveillance.

You may obtain other general information on your responsibilities under the Act from the Division of Small Manufacturers, International and Consumer Assistance at its toll-free number (800) 638-2041 or (301) 796-7100 or at its Internet address http://www.fda.gov/MedicalDevices/ResourcesforYou/Industry/default.htm.

Sincerely yours.

Anthony D. Watson, B.S., M.S., M.B.A.

Director

Division of Anesthesiology, General Hospital, Infection Control and Dental Devices Office of Device Evaluation Center for Devices and Radiological Health

INDICATIONS FOR USE

Applicant: SHANDONG JUI	NFENG INDUS	STRIES CO., LTD	
510(k) Number (if known): <u>*</u>	KISSS	ماما.	
Device Name: Powder Free Vin	yl Patient Exami	nation Gloves, Clear (non-colored)	
Indications For Use:		•	
Powder free vinyl patient example	mination glove i	s a disposable device intended for medica	ıl
purposes that is worn on the e	examiner's hand	or finger to prevent contamination betwee	n
patient and examiner.			
•			
		·	
Prescription Use(Part 21 CFR 801 Subpart D)	AND/OR	Over-The-Counter UseX_ (21 CFR 801 Subpart C)	
		•	
(PLEASE DO NOT WRITE BELOW	THIS LINE-CONT	INUE ON ANOTHER PAGE IF NEEDED)	
Conc	urrence of CDRH, (Office of Device Evaluation (ODE)	
	/ i.		

(Division Sign-Off)
Division of Anesthesiology, General Hospital
Infection Control, Dental Devices